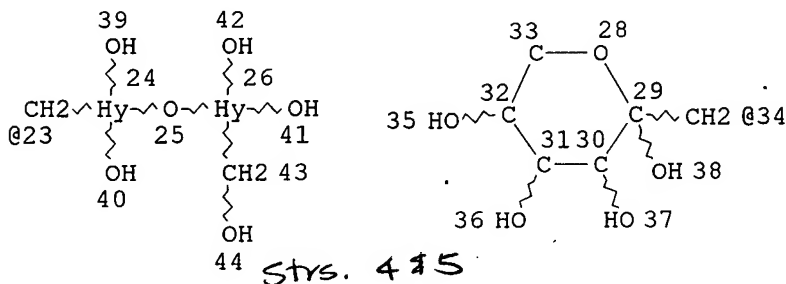
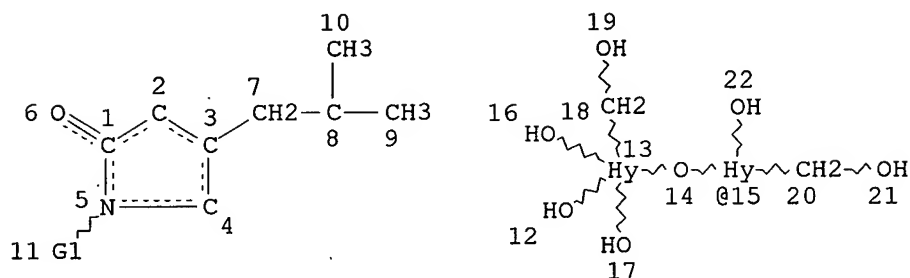


10/058903

(FILE 'REGISTRY' ENTERED AT 14:46:48 ON 12 JAN 2005)

L7

STR



VAR G1=15/23/34

NODE ATTRIBUTES:

CONNECT IS X2 RC AT 2

CONNECT IS X2 RC AT 4

CONNECT IS X2 RC AT 33

DEFAULT MLEVEL IS ATOM

GGCAT IS SAT AT 13

GGCAT IS SAT AT 15

GGCAT IS SAT AT 24

GGCAT IS SAT AT 26

DEFAULT ECLEVEL IS LIMITED

ECOUNT IS E1 O AT 13

ECOUNT IS E1 O AT 15

ECOUNT IS E1 O AT 24

ECOUNT IS E1 O AT 26

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 43

STEREO ATTRIBUTES: NONE

L9 12 SEA FILE=REGISTRY SSS FUL L7

100.0% PROCESSED 14218 ITERATIONS

12 ANSWERS

SEARCH TIME: 00.00.02

(FILE 'CAPLUS' ENTERED AT 15:07:11 ON 12 JAN 2005)

L10

2 S L9

L10 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:777769 CAPLUS

Searcher : Shears 571-272-2528

10/058903

DOCUMENT NUMBER: 137:284373  
 TITLE: Pregabalin-lactose conjugates for pharmaceuticals  
 INVENTOR(S): Hurley, Timothy Robert; Lovdahl, Michael James;  
 Tobias, Brian  
 PATENT ASSIGNEE(S): Warner-Lambert Company, USA  
 SOURCE: PCT Int. Appl., 36 pp.  
 CODEN: PIXXD2  
 DOCUMENT TYPE: Patent  
 LANGUAGE: English  
 FAMILY ACC. NUM. COUNT: 1  
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002078747	A2	20021010	WO 2002-IB647	20020225
WO 2002078747	A3	20031009		
WO 2002078747	C1	20031204		
WO 2002078747	C2	20031231		
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
US 2002187941	A1	20021212	US 2002-58903	20020128
CA 2440468	AA	20021010	CA 2002-2440468	20020225
EE 200300480	A	20031215	EE 2003-480	20020225
EP 1377318	A2	20040107	EP 2002-702614	20020225
R:	AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR			
BR 2002008439	A	20040323	BR 2002-8439	20020225
JP 2004524357	T2	20040812	JP 2002-577011	20020225
BG 108193	A	20040930	BG 2003-108193	20030924
NO 2003004348	A	20030929	NO 2003-4348	20030929
PRIORITY APPLN. INFO.:			US 2001-280176P	P 20010330
			WO 2002-IB647	W 20020225

AB Compns. containing pregabalin-lactose conjugates are useful for the treatment of of central nervous system disorders or diseases including seizure disorders, pain, depression, anxiety, sleep disorders, consumptive disorders, psychosis, tardive dyskinesia, Huntington's disease, or Parkinson's disease in humans. Pregabalin and lactose were dissolved in water and the solution was then heated at 90°. The resulting solid was then redissolved in approx. iso-PrOH by sonicating and heating. The product was subjected to reversed-phase preparative chromatog. to give (S)-1-[3,4-dihydroxy-6-hydroxymethyl-5-(3,4,5-trihydroxymethyl-6-hydroxymethyltetrahydropyran-2-yloxy)tetrahydropyran-2-yl]-4-isobutylpyrrolidin-2-one (I). Tablets contained I 25, lactose 50, corn starch (for mix) 10, corn starch (paste) 10, and Mg stearate 5 mg.

IT 466678-44-4P 466678-45-5P 466678-46-6P

466678-47-7P 466678-50-2P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological

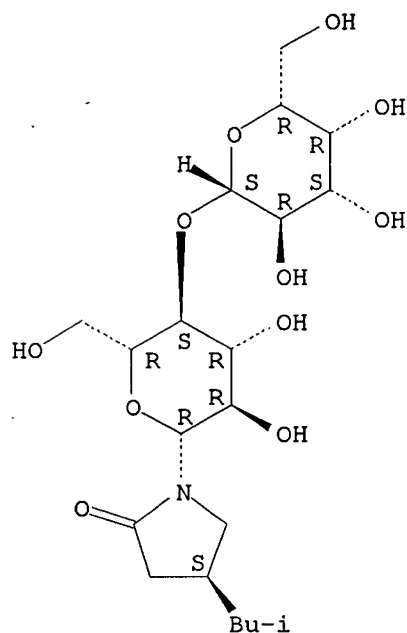
10/058903

study); PREP (Preparation); USES (Uses)  
(pregabalin lactose conjugates for pharmaceuticals)

RN 466678-44-4 CAPLUS

CN 2-Pyrrolidinone, 1-(4-O- $\beta$ -D-galactopyranosyl- $\beta$ -D-glucopyranosyl)-  
4-(2-methylpropyl)-, (4S)- (9CI) (CA INDEX NAME)

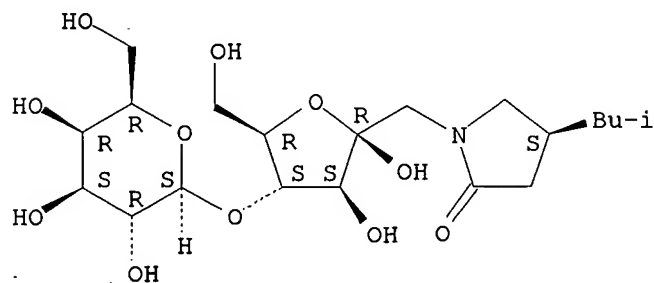
Absolute stereochemistry.



RN 466678-45-5 CAPLUS

CN  $\beta$ -D-Fructofuranose, 1-deoxy-4-O- $\beta$ -D-galactopyranosyl-1-[(4S)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

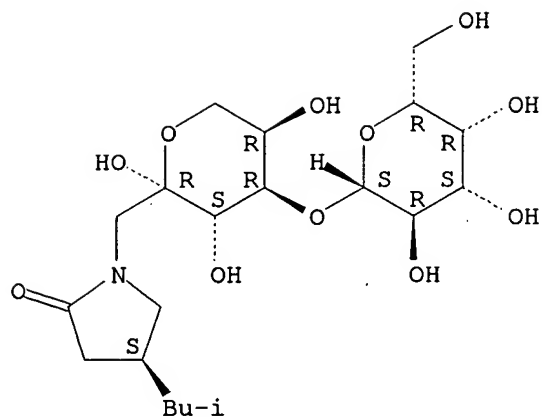


RN 466678-46-6 CAPLUS

CN  $\beta$ -D-Fructopyranose, 1-deoxy-4-O- $\beta$ -D-galactopyranosyl-1-[(4S)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

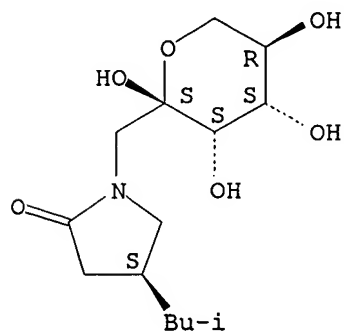
10/058903



RN 466678-47-7 CAPLUS

CN  $\alpha$ -D-Tagatopyranose, 1-deoxy-1-[(4S)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

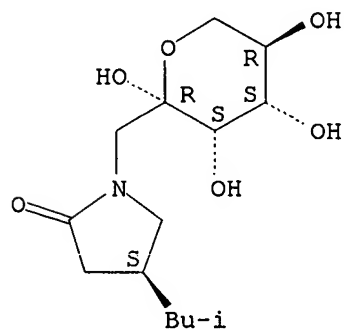
Absolute stereochemistry.



RN 466678-50-2 CAPLUS

CN  $\beta$ -D-Tagatopyranose, 1-deoxy-1-[(4S)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



Searcher :

Shears

571-272-2528

10/058903

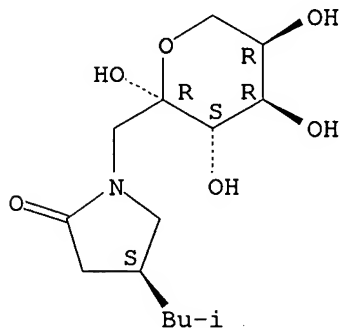
IT 466678-49-9

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)  
(pregabalin lactose conjugates for pharmaceuticals)

RN 466678-49-9 CAPLUS

CN  $\beta$ -D-Fructopyranose, 1-deoxy-1-[(4S)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L10 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:402961 CAPLUS

DOCUMENT NUMBER: 138:243397

TITLE: Synthesis and characterization of pregabalin lactose conjugate degradation products

AUTHOR(S): Lovdahl, Michael J.; Hurley, Timothy R.; Tobias, Brian; Priebe, Stephen R.

CORPORATE SOURCE: Analytical Development Department, Pfizer Global Research and Development, Ann Arbor, MI, 48105, USA

SOURCE: Journal of Pharmaceutical and Biomedical Analysis (2002), 28(5), 917-924

CODEN: JPBADA; ISSN: 0731-7085

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Seven degradation products observed in formulated pregabalin have been characterized. These compds. result from Maillard reactions and Amadori rearrangements. Heating pregabalin in the presence of lactose formed significant quantities of these degradation products. The seven compds. corresponding to the observed degradation products were isolated by

preparative

liquid chromatog. The synthesis, isolation, and spectral characterization of the degradation products are detailed.

IT 501665-97-0, PD 224377 501666-22-4, PD 310806

501666-23-5, PD 312237 501666-24-6, PD 312236

501666-25-7, PD 310886

RL: FMU (Formation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)

(synthesis and characterization of pregabalin lactose conjugate degradation products)

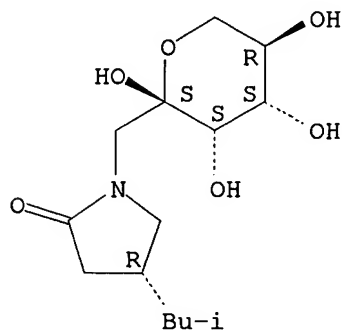
RN 501665-97-0 CAPLUS

Searcher : Shears 571-272-2528

10/058903

CN  $\alpha$ -D-Tagatopyranose, 1-deoxy-1-[(4R)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

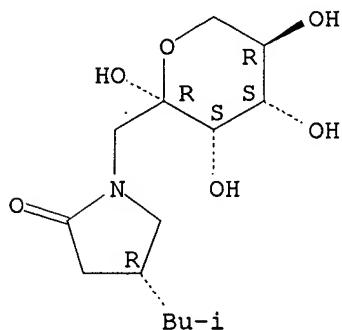
Absolute stereochemistry.



RN 501666-22-4 CAPLUS

CN  $\beta$ -D-Tagatopyranose, 1-deoxy-1-[(4R)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

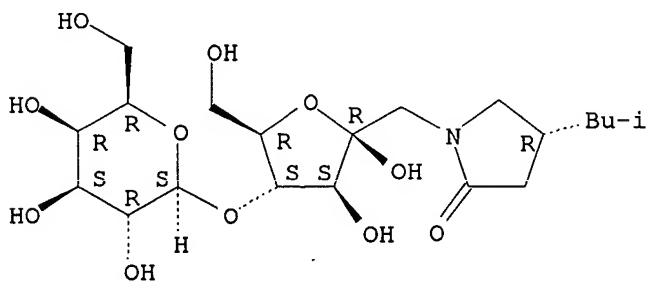
Absolute stereochemistry.



RN 501666-23-5 CAPLUS

CN  $\beta$ -D-Fructofuranose, 1-deoxy-4-O- $\beta$ -D-galactopyranosyl-1-[(4R)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

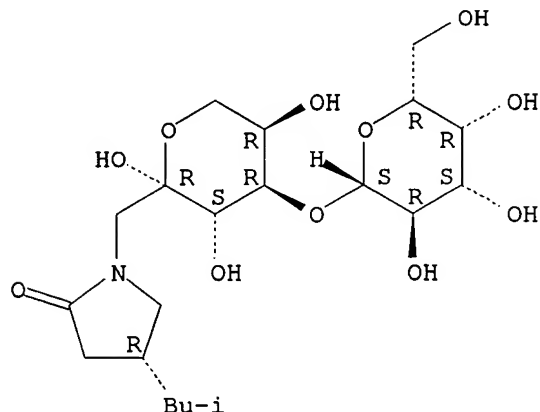


10/058903

RN 501666-24-6 CAPLUS

CN  $\beta$ -D-Fructopyranose, 1-deoxy-4-O- $\beta$ -D-galactopyranosyl-1-[(4R)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

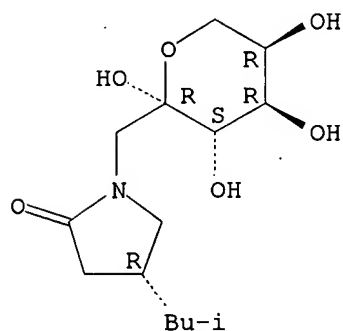
Absolute stereochemistry.



RN 501666-25-7 CAPLUS

CN  $\beta$ -D-Fructopyranose, 1-deoxy-1-[(4R)-4-(2-methylpropyl)-2-oxo-1-pyrrolidinyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



IT 501665-88-9, PD 224378

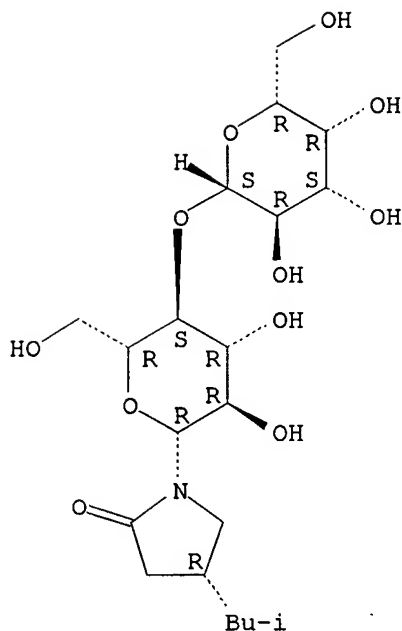
RL: FMU (Formation, unclassified); PRP (Properties); RCT (Reactant); FORM (Formation, nonpreparative); RACT (Reactant or reagent)  
(synthesis and characterization of pregabalin lactose conjugate degradation products)

RN 501665-88-9 CAPLUS

CN 2-Pyrrolidinone, 1-(4-O- $\beta$ -D-galactopyranosyl- $\beta$ -D-glucopyranosyl)-4-(2-methylpropyl)-, (4R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

10/058903



REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

FILE 'CAOLD' ENTERED AT 15:07:57 ON 12 JAN 2005  
L11 0 S L9

FILE 'USPATFULL' ENTERED AT 15:08:03 ON 12 JAN 2005  
L12 1 S L9

L12 ANSWER 1 OF 1 USPATFULL on STN

ACCESSION NUMBER: 2002:330258 USPATFULL

TITLE: Pregabalin lactose conjugates

INVENTOR(S): Hurley, Timothy Robert, Ann Arbor, MI, UNITED STATES  
Lovdahl, Michael James, Ann Arbor, MI, UNITED STATES  
Tobias, Brian, Ann Arbor, MI, UNITED STATES

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 2002187941	A1	20021212
APPLICATION INFO.:	US 2002-58903	A1	20020128 (10)

	NUMBER	DATE
PRIORITY INFORMATION:	US 2001-280176P	20010330 (60)
DOCUMENT TYPE:	Utility	
FILE SEGMENT:	APPLICATION	
LEGAL REPRESENTATIVE:	David R. Kurlandsky, Warner-Lambert Company, 2800 Plymouth Road, Ann Arbor, MI, 48105	
NUMBER OF CLAIMS:	7	
EXEMPLARY CLAIM:	1	
NUMBER OF DRAWINGS:	3 Drawing Page(s)	

*this applic*

Searcher : Shears 571-272-2528



LINE COUNT: 891

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB In accordance with the present invention, there is provided pregabalin lactose conjugate compounds.

Also provided as part of the present invention is a novel method of central nervous system disorders or diseases including seizure disorders, pain, depression, anxiety, sleep disorders, consumptive disorders, psychosis, tardive dyskinesia, Huntington's disease, or Parkinson's disease in a subject by administering to the subject a pharmaceutically effective amount of a pregabalin lactose conjugate.

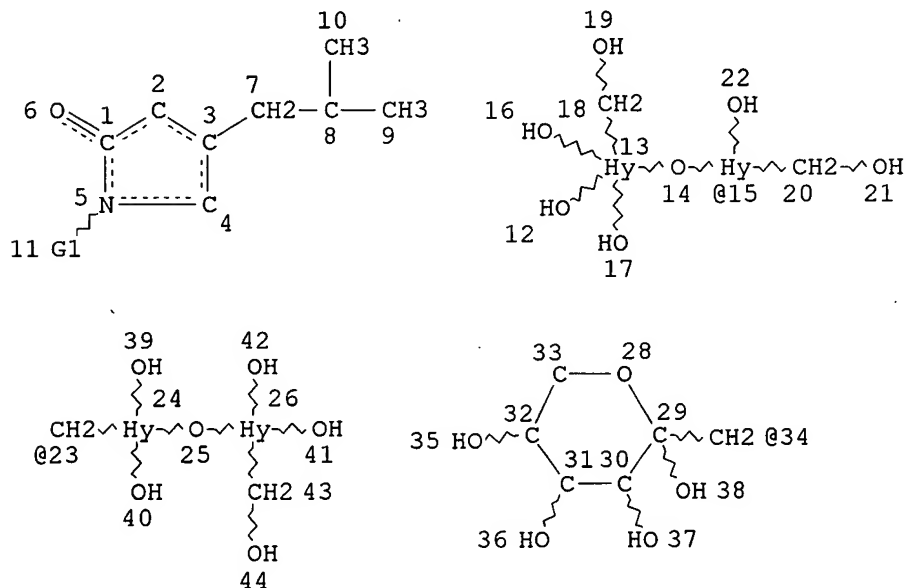
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

(FILE 'MEDLINE, BIOSIS, EMBASE' ENTERED AT 15:08:26 ON 12 JAN 2005)

L13 0 S L9

(FILE 'MARPAT' ENTERED AT 15:08:41 ON 12 JAN 2005)

L14 STR



VAR G1=15/23/34

NODE ATTRIBUTES:

CONNECT IS X2 RC AT 2

CONNECT IS X2 RC AT 4

CONNECT IS X2 RC AT 33

DEFAULT MLEVEL IS ATOM

MLEVEL IS CLASS AT 13 15 24 26

GGCAT IS SAT AT 13

GGCAT IS SAT AT 15

GGCAT IS SAT AT 24

GGCAT IS SAT AT 26

DEFAULT ECLEVEL IS LIMITED

ECOUNT IS E1 O AT 13

ECOUNT IS E1 O AT 15

10/058903

ECOUNT IS E1 O AT 24  
ECOUNT IS E1 O AT 26

GRAPH ATTRIBUTES:  
RSPEC I  
NUMBER OF NODES IS 43

STEREO ATTRIBUTES: NONE

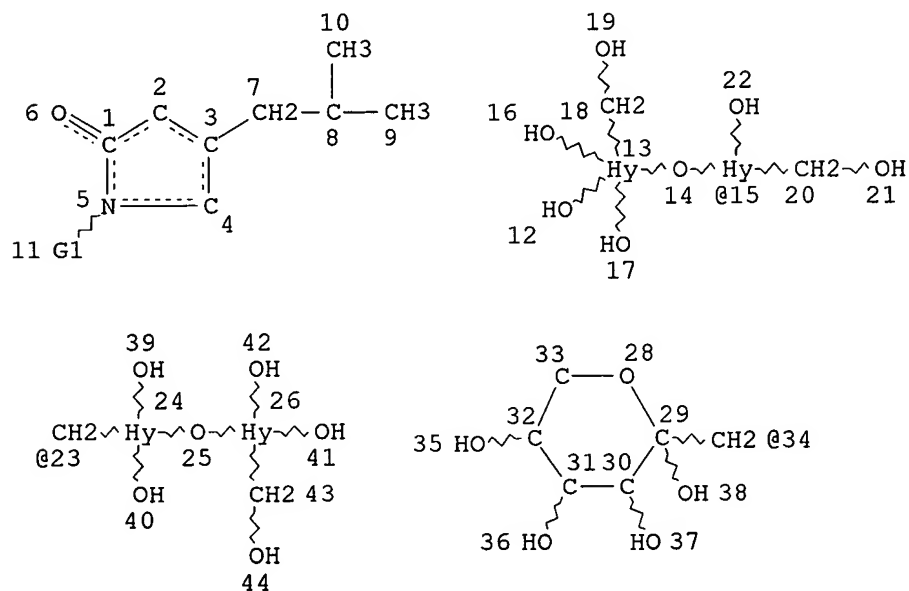
ATTRIBUTES SPECIFIED AT SEARCH-TIME:  
ECLEVEL IS LIM ON ALL NODES  
ALL RING(S) ARE ISOLATED

L16 0 SEA FILE=MARPAT SSS FUL L14 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 5780 ITERATIONS  
SEARCH TIME: 00.00.05

0 ANSWERS

FILE 'MARPATPREV' ENTERED AT 15:09:51 ON 12 JAN 2005  
L14 STR



VAR G1=15/23/34  
NODE ATTRIBUTES:  
CONNECT IS X2 RC AT 2  
CONNECT IS X2 RC AT 4  
CONNECT IS X2 RC AT 33  
DEFAULT MLEVEL IS ATOM  
MLEVEL IS CLASS AT 13 15 24 26  
GGCAT IS SAT AT 13  
GGCAT IS SAT AT 15  
GGCAT IS SAT AT 24  
GGCAT IS SAT AT 26  
DEFAULT ECLEVEL IS LIMITED  
ECOUNT IS E1 O AT 13

Searcher : Shears 571-272-2528

10/058903

ECOUNT IS E1 O AT 15  
ECOUNT IS E1 O AT 24  
ECOUNT IS E1 O AT 26

GRAPH ATTRIBUTES:  
RSPEC I  
NUMBER OF NODES IS 43

STEREO ATTRIBUTES: NONE

ATTRIBUTES SPECIFIED AT SEARCH-TIME:  
ECLEVEL IS LIM ON ALL NODES  
ALL RING(S) ARE ISOLATED

L17 0 SEA FILE=MARPATPREV SSS FUL L14 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 12 ITERATIONS 0 ANSWERS  
SEARCH TIME: 00.00.01

(FILE 'REGISTRY' ENTERED AT 15:13:22 ON 12 JAN 2005)  
L25 0 S ?"ISOBUTYL-1-(2,3,4,5-TETRAHYDROXY"?/CNS

L27 0 S ?"ISOBUTYL-1-(2,3,5-TRIHIDROXY"?/CNS  
L28 0 S ?"ISOBUTYL-1-(2,3,4-TRIHIDROXY"?/CNS

- Named  
compds.

L31 0 S ?"DIHYDROXY-6-HYDROXYMETHYL-5-(3,4,5-TRIHIDROXY"?/CNS  
L32 0 S ?"DIHYDROXY-5-HYDROXYMETHYL-4-(3,4,5-TRIHIDROXY"?/CNS

(FILE 'CAPLUS' ENTERED AT 15:20:41 ON 12 JAN 2005)  
L33 1381 SEA FILE=CAPLUS ABB=ON PLU=ON (DIHYDROXY OR DI HYDROXY) (S) (HY  
DROXYMETHYL OR HYDROXY (W) (ME OR METHYL))  
L34 128 SEA FILE=CAPLUS ABB=ON PLU=ON L33 (S) (TETRAHYDRO? OR TETRA  
HYDRO?)  
L35 1 SEA FILE=CAPLUS ABB=ON PLU=ON L34 (S) (ISOBUTYL? OR (ISO OR  
I) (W) (BUTYL?))

L36 692 SEA FILE=CAPLUS ABB=ON PLU=ON 4 (W) (ISOBUTYL OR (ISO OR  
I) (W) (BUTYL OR BU))  
L37 0 SEA FILE=CAPLUS ABB=ON PLU=ON L36 (S) (4 (W) 5 (W) (TETRAHYDROXY?  
OR TETRA HYDROXY?))

L36 692 SEA FILE=CAPLUS ABB=ON PLU=ON 4 (W) (ISOBUTYL OR (ISO OR  
I) (W) (BUTYL OR BU))  
L38 0 SEA FILE=CAPLUS ABB=ON PLU=ON L36 (S) (5 (W) (TRIHIDROXY? OR TRI  
HYDROXY?))

L39 0 L35 NOT L10

(FILE 'MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS,  
JAPIO' ENTERED AT 15:25:06 ON 12 JAN 2005)  
L40 1 S L35

Searcher : Shears 571-272-2528

10/058903

L41 0 S L37  
L42 0 S L38

L40 ANSWER 1 OF 1 WPIDS COPYRIGHT 2005 THE THOMSON CORP on STN  
ACCESSION NUMBER: 2002-241176 [29] WPIDS  
DOC. NO. CPI: C2002-072446  
TITLE: New beta-hydroxyamide compound useful in powder coating  
compositions as crosslinkers and/or curing agents.  
DERWENT CLASS: A25 A60 E16 G02  
INVENTOR(S): MANEA, M; PETERSSON, C  
PATENT ASSIGNEE(S): (PEST) PERSTORP SPECIALTY CHEM AB; (MANE-I) MANEA M;  
(PETE-I) PETERSSON C  
COUNTRY COUNT: 95  
PATENT INFORMATION:

PATENT NO	KIND	DATE	WEEK	LA	PG
WO 2001098257	A1	20011227	(200229)*	EN	23
RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TR TZ UG ZW					
W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW					
AU 2001074750	A	20020102	(200230)		
EP 1292567	A1	20030319	(200322)	EN	
R: AL AT BE CH CY DE DK ES FI FR GB GR IE IT LI LT LU LV MC MK NL PT RO SE SI TR					
US 2003195373	A1	20031016	(200369)		
JP 2004501133	W	20040115	(200410)		42

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 2001098257	A1	WO 2001-SE1359	20010615
AU 2001074750	A	AU 2001-74750	20010615
EP 1292567	A1	EP 2001-941393	20010615
		WO 2001-SE1359	20010615
US 2003195373	A1	WO 2001-SE1359	20010615
		US 2003-311295	20030204
JP 2004501133	W	WO 2001-SE1359	20010615
		JP 2002-504213	20010615

FILING DETAILS:

PATENT NO	KIND	PATENT NO
AU 2001074750	A Based on	WO 2001098257
EP 1292567	A1 Based on	WO 2001098257
JP 2004501133	W Based on	WO 2001098257

PRIORITY APPLN. INFO: SE 2000-2268 20000619  
AN 2002-241176 [29] WPIDS  
AB WO 200198257 A UPAB: 20020508  
NOVELTY - A beta -hydroxyamide compound (I) is new.

Searcher : Shears 571-272-2528

10/058903

DETAILED DESCRIPTION - A beta -hydroxyamide compound of formula (I) is new.

R1 = alkyl, alkoxyalkyl, hydroxyalkyl or hydroxyalkoxyalkyl;

R2 = alkyl, aryl, alkylaryl or arylalkyl;

R3 = N-alkyl or N-cycloalkyl having at least one OH in beta-position;

m, n = 1 or more.

An INDEPENDENT CLAIM is also included for a process for the synthesis of (I).

USE - In powder coating compositions as chemical intermediates, crosslinkers and/or curing agents.

ADVANTAGE - (I) provides a coating that can be formulated as a solvent-borne or water-borne system, and solvent-borne coatings can be formulated as high-solid systems. It allows crosslinking temperature to be moderate-to-high in the range of 150-200 deg. C, and reduces the compatibility problems due to di, tri or polyhydric core compounds. The coating obtained is clear and has improved flexibility and hardness.  
Dwg.0/0

FILE 'HOME' ENTERED AT 15:28:43 ON 12 JAN 2005